=> fil hcaplu
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FILE COVERS 1907 - 1 Feb 2002 VOL 136 ISS 6 FILE LAST UPDATED: 30 Jan 2002 (20020130/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

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The P indicator for Preparations was not generated for all of the CAS Registry Numbers that were added to the CAS files between 12/27/01 and 1/23/02. As of 1/23/02, the situation has been resolved. Searches and/or SDIs in the H/Z/CA/CAplus files incorporating CAS Registry Numbers with the P indicator executed between 12/27/01 and 1/23/02 may be incomplete. See the NEWS message on this topic for more information.

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=> d stat que
             53 SEA FILE=REGISTRY SERUM ALBUMIN?/CN
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L3
             63 SEA FILE=REGISTRY ANGIOSTA?/CN
L4
             42 SEA FILE=REGISTRY ENDOSTATIN?
L5
            342 SEA FILE=REGISTRY CYSTINE?/CN
L7
          61673 SEA FILE=HCAPLUS L1 OR SERUM(W) ALBUMIN OR SA
L13
          12190 SEA FILE=HCAPLUS CHIMERIC (5A) (PROTEIN? OR ?PEPTIDE?) OR L3
L14
            590 SEA FILE=HCAPLUS L4 OR ANGIOSTA?
L15
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L1'9
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              9 SEA FILE=HCAPLUS L20 AND L14
L21
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=> d ibib abs hitrn 121 1-9

Davis 09/768,183 Page 2

L21 ANSWER 1 OF 9 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:936090 HCAPLUS

DOCUMENT NUMBER:

136:58776

TITLE:

Chimeric polypeptides of

serum albumin and uses related

thereto

INVENTOR(S):

Gyuris, Jeno; Lamphere, Lou

PATENT ASSIGNEE(S):

SOURCE:

USA U.S. Pat. Appl. Publ., 34 pp., Cont.-in-part of U.S.

Ser. No. 619,285.

CODEN: USXXCO

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

APPLICATION NO. DATE KIND DATE PATENT NO. US 2001-764918 20010118 US 2001056075 A1 20011227 US 1999-144534 P 19990719 PRIORITY APPLN. INFO.: US 2000-619285 A2 20000719

The present invention relates to chimeric polypeptides AB in which a serum albumin protein has been altered to include one or more biol. active heterologous peptide sequences. The chimeric polypeptides may exhibit therapeutic activity related to the heterologous peptide sequences coupled with the improved serum half-lives derived from the serum albumin protein fragments. Heterologous peptide sequences may be chosen to promote any biol. effect, including angiogenesis inhibition, antitumor activity, and induction of apoptosis. The therapeutic effect may be achieved by direct administration of the chimeric polypeptide, or by transfecting cells with a vector including a nucleic acid encoding such a chimeric polypeptide.

IT 86090-08-6P, Angiostatin 187888-07-9P,

Endostatin

RL: BPN (Biosynthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(fusion proteins; chimeric polypeptides of serum albumin and uses related thereto)

L21 ANSWER 2 OF 9 HCAPLUS COPYRIGHT 2002 ACS 2001:781079 HCAPLUS

ACCESSION NUMBER: DOCUMENT NUMBER:

135:348851

TITLE:

Albumin fusion proteins with therapeutic proteins for

improved shelf-life

INVENTOR(S):

Rosen, Craig A.; Haseltine, William A.

Human Genome Sciences, Inc, USA PATENT ASSIGNEE(S):

SOURCE:

PCT Int. Appl., 606 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

sequences. The chimeric polypeptides may exhibit therapeutic activity related to the heterologous peptide sequences coupled with the improved serum half-lives derived from the serum albumin protein fragments. Heterologous peptide sequences may be chosen to promote any biol. effect, including angiogenesis inhibition, antitumor activity, and induction of apoptosis. The therapeutic effect may be achieved by direct administration of the chimeric polypeptide, or by transfecting cells with a vector including a nucleic acid encoding such a chimeric polypeptide. 86090-08-6P, Angiostatin 187888-07-9P, Endostatin

IT

RL: BPN (Biosynthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(fusion proteins; chimeric polypeptides of serum albumin and uses related thereto)

L21 ANSWER 2 OF 9 HCAPLUS COPYRIGHT 2002 ACS 2001:781079 HCAPLUS

ACCESSION NUMBER:

DOCUMENT NUMBER:

135:348851

TITLE:

Albumin fusion proteins with therapeutic proteins for

improved shelf-life

INVENTOR(S):

Rosen, Craig A.; Haseltine, William A.

Human Genome Sciences, Inc, USA

PATENT ASSIGNEE(S): SOURCE:

PCT Int. Appl., 606 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English 7

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PAT | PATENT NO. | | | | KIND DATE | | | | A | PPLI | CATI | DATE | | | | | | |
|----------|------------------|-----|-----|-------------|-----------|------|-----|-----------------------|----------------|------|------|------|------------|------|------|-----|-----|--|
| WO | 2001079444 | | | A2 20011025 | | 1025 | | WO 2001-US12013 | | | | | 3 20010412 | | | | | |
| | W: AE, AG, | | | AL, | AM, | AT, | AU, | ΑZ, | BA, | BB, | BG, | BR, | BY, | ΒZ, | CA, | CH, | CN, | |
| | | CO, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ, | EE, | ES, | FI, | GB, | GD, | GE, | GH, | GM, | |
| | | HR, | HU, | ID, | IL, | IN, | IS, | JP, | ΚE, | KG, | ΚP, | KR, | ΚZ, | LC, | LK, | LR, | LS, | |
| | | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | ΜX, | ΜZ, | NO, | ΝZ, | PL, | PT, | RO, | |
| | | RU, | SD, | SE, | SG, | SI, | SK, | SL, | ТJ, | TM, | TR, | TT, | ΤZ, | UΑ, | UG, | US, | UZ, | |
| | | VN, | YU, | ZA, | ZW, | AM, | ΑZ, | BY, | KG, | ΚZ, | MD, | RU, | ТJ, | TM | | | | |
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| | | DE, | DK, | ES, | FI, | FR, | GB, | GR, | ΙE, | IT, | LU, | MC, | ΝL, | PT, | SE, | TR, | BF, | |
| | | ВJ, | CF, | CG, | CI, | CM, | GA, | GN, | GW, | ML, | MR, | ΝE, | SN, | TD, | TG | | | |
| PRIORITY | TY APPLN. INFO.: | | | | | | | | US 2000-229358 | | | | P 20000412 | | | | | |
| | • | | | | | | | US 2000-199384 P 2000 | | | | | 2000 | 0425 | | | | |
| | | | | | | | | | US 2 | -000 | 2569 | 31 | P | 2000 | 1221 | | | |

The present invention encompasses fusion proteins of albumin with various AΒ therapeutic proteins. Therapeutic proteins may be stabilized to extend the shelf-life, and/or to retain the therapeutic protein's activity for extended periods of time in soln., in vitro and/or in vivo, by genetically or chem. fusing or conjugating the therapeutic protein to albumin or a fragment or variant of albumin. Use of albumin fusion proteins may also reduce the need to formulate the protein solns. with large excesses of carrier proteins to prevent loss of therapeutic proteins due to factors

such as binding to the container. Nucleic acid mols. encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors contg. these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Thus, plasmid vectors are constructed in which DNA encoding the desired therapeutic protein may be inserted for expression of the albumin fusion proteins in yeast (pPPC0005) and mammalian cells (pC4:HSA). Yeast-derived signal sequences from Saccharomyces cerevisiae invertase SUC2 gene, or the stanniocalcin or native human serum albumin signal peptides, are used for secretion in yeast or mammalian systems, resp. Thus, the fusion product of human growth hormone with residues 1-387 of human serum albumin retains essentially intact biol. activity after 5 wk of incubation in tissue culture media at 37.degree., whereas recombinant human growth hormone used as control lost its biol. activity in the first week. Although the potency of the albumin fusion proteins is slightly lower than the unfused counterparts in rapid bioassays, their biol. stability results in much higher biol. activity in the longer term in vitro assay or in vivo assays. Addnl., the present invention encompasses pharmaceutical compns. comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disorders or conditions using albumin fusion proteins of the invention.

IT 187888-07-9P, Endostatin

RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (albumin fusion proteins with therapeutic proteins for improved shelf-life)

L21 ANSWER 3 OF 9 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:781078 HCAPLUS

DOCUMENT NUMBER:

135:348850

TITLE:

Albumin fusion proteins with therapeutic proteins for

improved shelf-life

INVENTOR(S):

Rosen, Craig A.; Haseltine, William A. Human Genome Sciences, Inc., USA

PATENT ASSIGNEE(S):

PCT Int. Appl., 374 pp.

CODEN. PIYY

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

. 7

FAMILY ACC. NUM. COUNT: 7

PATENT INFORMATION:

| PATENT | KIND DATE | | | | A | PPLI | CATI | o. : | DATE | | | | | | | |
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| | | | | | | | | | | | | | | | | |
| WO 2001 | 43 | A2 20011025 | | | | WO 2001-US11924 20010412 | | | | | | | | | | |
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| | co. | CR. | CU. | CZ, | DE, | DK, | DM, | DZ, | EE, | ES, | FI, | GB, | GD, | GE, | GH, | GM, |
| | HR. | HU. | ID. | IL. | IN, | IS, | JP, | KE, | KG, | KP, | KR, | KZ, | LC, | LK, | LR, | LS, |
| • | LT. | LU. | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NO, | NZ, | PL, | PT, | RO, |
| | RU. | SD. | SE. | SG, | SI, | SK, | SL, | TJ, | TM, | TR, | TT, | TZ, | UA, | UG, | US, | UΖ, |
| | VN. | YU. | ZA. | ZW. | AM, | AZ, | BY, | KG, | KZ, | MD, | RU, | TJ, | TM | | | |
| ₽W• | GH. | GM. | KE. | LS. | MW. | MZ, | SD, | SL, | SZ, | TZ, | UG, | ZW, | AT, | BE, | CH, | CY, |
| ***** | DE, | DK, | ES, | FI, | FR, | GB, | GR, | IE, | IT, | LU, | MC, | NL, | PT, | SE, | TR, | BF, |

BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG US 2000-229358 P 20000412 PRIORITY APPLN. INFO.: US 2000-199384 P 20000425 US 2000-256931 Р 20001221

The present invention encompasses fusion proteins of albumin with various AΒ therapeutic proteins. Therapeutic proteins may be stabilized to extend the shelf-life, and/or to retain the therapeutic protein's activity for extended periods of time in soln., in vitro and/or in vivo, by genetically or chem. fusing or conjugating the therapeutic protein to albumin or a fragment or variant of albumin. Use of albumin fusion proteins may also reduce the need to formulate the protein solns. with large excesses of carrier proteins to prevent loss of therapeutic proteins due to factors such as binding to the container. Nucleic acid mols. encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors contg. these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Thus, plasmid vectors are constructed in which DNA encoding the desired therapeutic protein may be inserted for expression of the albumin fusion proteins in yeast (pPPC0005) and mammalian cells (pC4:HSA). Yeast-derived signal sequences from Saccharomyces cerevisiae invertase SUC2 gene, or the stanniocalcin or native human serum albumin signal peptides, are used for secretion in yeast or mammalian systems, resp. Thus, the fusion product of human growth hormone with residues 1-387 of human serum albumin retains essentially intact biol. activity after 5 wk of incubation in tissue culture media at 37.degree., whereas recombinant human growth hormone used as control lost its biol. activity in the first week. Although the potency of the albumin fusion proteins is slightly lower than the unfused counterparts in rapid bioassays, their biol. stability results in much higher biol. activity in the longer term in vitro assay or in vivo assays. Addnl., the present invention encompasses pharmadeutical compns. comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disorders or conditions using albumin fusion proteins of the invention.

86090-08-6P, Angiostatin IT

RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (albumin fusion proteins with therapeutic proteins for improved shelf-life)

L21 ANSWER 4 OF 9 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: DOCUMENT NUMBER:

2001:64027 HCAPLUS

134:110478

TITLE:

Chimeric polypeptides of serum albumin containing

heterologous peptide sequences, and

therapeutic uses thereof Gyuris, Jeno; Lamphere, Lou

INVENTOR(S):

GPC Biotech Inc., USA.

PATENT ASSIGNEE(S): SOURCE:

PCT Int. Appl., 45 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 2 PATENT INFORMATION:

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APPLICATION NO.
                     KIND DATE
    PATENT NO.
                                           WO 2000-US19689 20000719
    WO 2001005826
                      A2
                            20010125
                            20010802
    WO 2001005826
                     A3
            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
            HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
            LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
            SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU,
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        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
             CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                        US 1999-144534
PRIORITY APPLN. INFO .:
    The invention discloses chimeric polypeptides in which
     a serum albumin protein has been altered to include
     one or more biol. active heterologous peptide
     sequences. The chimeric polypeptides may exhibit
     therapeutic activity related to the heterologous peptide
     sequences coupled with the improved serum half-lives derived from the
     serum albumin protein fragments. Heterologous
    peptide sequences may be chosen to promote any biol. effect,
     including angiogenesis inhibition, antitumor activity, and induction of
     apoptosis. The therapeutic effect may be achieved by direct
     administration of the chimeric polypeptide, or by
     transfecting cells with a vector including a nucleic acid encoding such a
     chimeric polypeptide.
     86090-08-6D, Angiostatin, fragments 187888-07-9D
IT
     , Endostatin, fragments
     RL: BAC (Biological activity or effector, except adverse); BOC (Biological
     occurrence); THU (Therapeutic use); BIOL (Biological study); OCCU
     (Occurrence); USES (Uses)
        (chimeric polypeptides of serum
        albumin contg. heterologous peptide
        sequences, and therapeutic uses thereof)
L21 ANSWER 5 OF 9 HCAPLUS COPYRIGHT 2002 ACS
                         1998:175372 HCAPLUS
ACCESSION NUMBER:
                         128:229741
DOCUMENT NUMBER:
                         Methods of enhancing production performance of birds
TITLE:
                         comprising administration of heterologous
                         protein comprising avian .alpha.-subunit
                         inhibin protein
                         Fioretti, William C.; Kousoulas, Konstantin;
INVENTOR(S):
                         Satterlee, Daniel G.
                         Agritech Technologies, Ltd., USA; Board of Supervisors
PATENT ASSIGNEE(S):
                         of Louisiana State Univ. and Agricultural & Mechanical
                         College
                         U.S., 29 pp. Cont.-in-part of U.S. Ser. No. 395,554,
SOURCE:
                         abandoned.
                         CODEN: USXXAM
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Page 7 Davis ,09/768,183

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| | PATENT NO. | | | | | KIND DAT | | | | A: | PPLI | CATI | ON NO | o. | DATE | | | |
|----|------------|-------|-------------------|------|-------------|----------|----------|------|-----|------------------------|------|------|-----------|--------|------|------|-----|-----|
| | US | 5725 | - - 858 | | A | | 1998 | 0310 | | US 1995-481633 1995060 | | | | | | | | |
| | US | 5747 | 659 | | A | | 1998 | 0505 | | U | S 19 | 95-4 | 80493 | 3 | 1995 | 0607 | • | |
| | CA | 2222 | 947 | | A | A. | 19961219 | | | C | A 19 | 96-2 | 2229 | 47 | 1996 | 0606 | | |
| | WO | 9640 | 219 | | A1 19961219 | | | | W | O 19 | 96-U | 9 | 1,9960606 | | | | | |
| | | w: | AL, | AM, | ΑT, | AU, | AZ, | BB, | BG, | BR, | BY, | CA, | CH, | CN, | CZ, | DE, | DK, | EE, |
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| | | | SE, | SG | | | | | | | | | | | | | | |
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| | AU | 9660 | 941 | | Α | 1 | 1996 | 1230 | | A ¹ | U 19 | 96-6 | 0941 | | 1996 | 0606 | | |
| | ΑU | 7263 | 21 | | B | 2 | 2000 | 1102 | | | | | | | | | | |
| | EP | 8336 | 58 | | Α | 1 | 1998 | 0408 | | E | P 19 | 96-9 | 1823 | 4 | 1996 | 0606 | | |
| | | R: | ΑT, | BE, | CH, | DE, | DK, | ES, | FR, | GB, | GR, | IT, | LI, | LU, | ΝL, | SE, | MC, | PT, |
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| | CN | 1192 | 692 | | Α | | 1998 | 0909 | | C | N 19 | 96-1 | 9583 | 9 | 1996 | 0606 | | |
| | JP | 1151 | 1964 | | T | 2 | 1999 | 1019 | | _ | | | | | 1996 | | | |
| PR | IORIT | Y APP | LN. | INFO | . : | | | | | | | | | | 1994 | | | |
| | | | | | | | | | | | | | | | 1995 | | | |
| | | | | | | | | | • | US 1 | 995- | 4816 | 33 | Α3 | 1995 | 0607 | | |
| | | | | | | | | | 1 | WO 1 | 996- | US92 | 29 | W | 1996 | 0606 | | |

The prodn. performance of avians is enhanced by administering a AΒ heterologous fusion protein comprised of inhibin, or a fragment thereof, and a carrier protein. A DNA fragment (cINA521) was excised from the chicken inhibit .alpha.-subunit cDNA clone cINA6 using PstI digestion and cloned in the com. plasmid vector pMAL-c in .frame with the maltose-binding protein (MBP) and a fusion protein of appropriate size was detected after IPTG induction and SDS-PAGE. The resulting protein conjugate (MBP-cINA521) was used as an antigen to immunize pre-pubescent, female Japanese quail (Coturnix coturnix japonica) against circulating inhibin levels. MBP-cINA521 immunization enhances prodn. performance as it accelerates the onset of puberty, increases egg lay intensity, and accelerates the onset of max. egg lay in Japanese quail. Improved prodn. performance is also obsd. in ostrich, emu, chicken, turkey, and parrots.

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L21 ANSWER 6 OF 9 HCAPLUS COPYRIGHT 2002 ACS
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ACCESSION NUMBER:

1997:679116 HCAPLUS

DOCUMENT NUMBER:

127:330381

TITLE:

Modified/chimeric superantigens and their use

Antonsson, Per; Hansson, Johan; Bjork, Per; Dohlsten, INVENTOR(S):

Mikael; Kalland, Terje; Abrahmsen, Lars; Forsberg,

Goran

PATENT ASSIGNEE(S):

Pharmacia & Upjohn AB, Swed.; Antonsson, Per; Hansson, Johan; Bjork, Per; Dohlsten, Mikael; Kalland, Terje;

Abrahmsen, Lars; Forsberg, Goran

PCT Int. Appl., 58 pp. . SOURCE:

CODEN: PIXXD2

Davis ,09/768,183 Page 8

DOCUMENT TYPE:

Patent English

LANGUAGE:

1 1

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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APPLICATION NO.
                     KIND DATE
    PATENT NO.
                                          WO 1997-SE537
                                                            19970326
                           19971009
                      A1
    WO 9736932
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            LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT,
            RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN,
            AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
        RW: GH, KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB,
            GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN,
            ML, MR, NE, SN, TD, TG
                                           CA 1997-2222757 19970326
                           19971009
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                                           AU 1997-25251
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                            19990722
    AU 707827
                      В2
                                                            19970326
                                           EP 1997-916693
                            19980415
     EP 835266
                      A1
           AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, FI
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                            19980129
     NO 9705435
                                                         A 19960329
PRIORITY APPLN. INFO.:
                                        SE 1996-1245
                                                         A 19960812
                                        US 1996-695692
                                                        W 19970326
                                        WO 1997-SE537
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A conjugate between a target-seeking moiety and a modified superantigen, ΑB characterized in that the superantigen is a wild-type superantigen (SA I) in which an amino acid residue in a superantigen region (region I) detg. binding to TCR, preferably TCR V.beta., and T cell activation have been replaced by another amino acid residue while retaining the ability to activate a subset of T cells. In a preferred embodiment the modified superantigen is a chimer between at least two wild-type superantigens (SA I, SA II etc.) characterized in that one or more amino acid residues in a region detg. binding to TCR and T cell activation have been interchanged between various wild-type superantigens. A therapeutic method making use of modified/chimeric superantigens as defined in the preceding paragraphs. An antibody prepn. in which the cysteine residues that provide for interchain disulfide bonds have been mutated so as to forbid interchain disulfide bridges, preferably to serine residues, for use as a pharmaceutical. Plasmid 5T4Fab-SEA encoding fusion protein contg. antibody 5T4 variable region and murine IgG1 V.kappa. chain and Staphylococcal enterotoxin A was constructed, and the expressed chimeric superantigen was tested.

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L21 ANSWER 7 OF 9 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1996:617214 HCAPLUS
```

DOCUMENT NUMBER: 126:3980

TITLE:

Anomalous mobility of sulfitolyzed proteins in

SDS-PAGE. Analysis and applications

.09/768,183 Page 9 Davis

AUTHOR(S):

Malhotra, M.; Sahal, D.

CORPORATE SOURCE:

International Centre Genetic Eng. Biotechnology, Aruna

Asaf Ali Marg, Recombinant Gene Products, New Delhi,

India

SOURCE:

Int. J. Pept. Protein Res. (1996), 48(3), 240-248

CODEN: IJPPC3; ISSN: 0367-8377

PUBLISHER: DOCUMENT TYPE: Munksgaard Journal

LANGUAGE:

English

Some cysteine-contg. proteins upon sulfitolysis have been found AΒ to show anomalously retarded SDS-PAGE mobilities in non-reducing gels. These proteins include bovine serum albumin, ovalbumin, aldolase, RNase and a recombinant fusion protein (XA)

consisting of a portion of .gamma.-interferon linked to the A chain of human insulin. This mobility shift has been employed to det. the stability of the sulfonated products and to study the kinetics of the sulfitolysis reaction. Partially sulfonated products of intermediate shifts were obsd. at 0.01% .beta.-mercaptoethanol (.beta.-ME), while 0.05% .beta.-ME gave a shift characteristic of the completely reduced protein. The undiluted sulfitolysis reagent reacted with XA to give within 1 min a gel shift characteristic of the fully sulfitolyzed protein. Its transition stages could be visualized at 15, 30 and 60 min when the reagent was dild. four-fold. In the presence of 8 M urea, the sulfitolysis of BSA was nearly complete at 30 min when the sulfitolysis reagent was used at a diln. of 1:5. However, under the same conditions BSA was predominantly unsulfitolyzed in the absence of urea. In order to elucidate the mechanism of sulfonation shift, several derivs. of XA, e.g. performic acid oxidized, alkylated with (a) iodoacetamide and (b) iodoacetate, have been prepd. While the mobility of XASSO3- was sensitive to the presence of .beta.-ME, all other derivs. moved in a .beta.-ME-insensitive fashion. Furthermore, while the nonreducing mobilities of the acidic derivs. (-SSO3-, -SO3- and -SCH2CO2-) were anomalously retarded and identical, the mobility of the iodoacetamide deriv. was intermediate between the retarded acidic derivs. above and XA below. These studies have suggested a role of the extended conformation of the A chain of insulin in causing a mobility shift of the acidic derivs. in this series. Similar results were obsd. in an analogous series of derivs. prepd. from BSA. Non-denaturing gel filtration analyses of native vs. sulfitolyzed samples of serum albumin, ovalbumin and RNase have indicated that the sulfitolyzed proteins elute earlier than their native counterparts and appear to be significantly larger than their true mol. wts. CD anal. has indicated significant loss in helicity of sulfitolyzed BSA. This suggests that the retarded mobility of sulfitolyzed proteins seen on SDS-PAGE is likely to be due to an expansion in the hydrodynamic vols. of these proteins, a phenomenon triggered by cleavage of disulfide bonds and further accentuated by the

L21 ANSWER 8 OF 9 HCAPLUS COPYRIGHT 2002 ACS 1991:507784 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

115:107784

introduction of strongly neg. charged sulfonates.

TITLE:

Production of heterologous proteins

in plants or plant cells by recombinant DNA techniques

Mogen International N. V., Neth. . PATENT ASSIGNEE(S):

Page 10 Davis .09/768,183

SOURCE:

Neth. Appl., 37 pp.

CODEN: NAXXAN

DOCUMENT TYPE:

Patent Dutch

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|-------------------------|------------|-----------|-------------------------------|----------------------|
| NL 8901932 | A | 19910218 | NL 1989-1932 WO 1990-NL108 | 19890726 19900726 |
| WO 9102066 W: JP, US | A1 | 19910221 | | 19900726 |
| RW: AT, BE, | CH, DE | , DK, ES, | FR, GB, IT, LU, NL, SE | |
| EP 436003 | A1 | 19910710 | EP 1990-911488 | 19900726 |
| R: AT, BE, | CH, DE | , DK, ES, | FR, GB, IT, LI, LU, NL, | SE |
| JP 04502861 | T 2 | 19920528 | JP 1990-510940 | 19900726 |
| US 5716802 | Α | 19980210 | US 1991-659287 | 19910521 |
| | | 19970722 | US 1995-469856 | 19950606 |
| US 5763748 | A | 19980609 | US 1997-829057 | 19970331 |
| PRIORITY APPLN. INFO | . : | | NL 1989-1932 | 19890726 |
| | | | WO 1990-NL108 | 19900726 |
| | | | US 1991-659287 | 19910521 |

Prodn. and secretion of a heterologous protein in a AB plant or plant cell is made possible by substituting a signal sequence functional in the plant host for the natural signal sequence immediately preceding the protein gene in the expression cassette introduced into the plant host. Thus, cloned cDNA for human prepro-serum albumin was manipulated to insert a sequence coding for a signal peptide from Samsun NN tobacco protein PROB12 immediately prior to the sequence coding for mature human serum albumin. Plasmid pMOG236, an Agrobacterium tumefaciens binary vector contg. this construct, was used to transform potato slices which were regenerated to mature transgenic plants. Leaves, stems, and tubers of these plants contained human serum albumin. The albumin was secreted by leaf cells and was found at elevated concn. in the extracellular fluid.

L21 ANSWER 9 OF 9 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1991:201173 HCAPLUS

DOCUMENT NUMBER:

114:201173

TITLE:

Albumin fragment gene and its cloning and expression

in Escherichia coli

INVENTOR(S):

Maki, Noboru; Yagi, Shintaro; Suzuki, Masanori

PATENT ASSIGNEE(S):

SOURCE:

Toa Nenryo Kogyo K. K., Japan

Jpn. Kokai Tokkyo Koho, 24 pp. CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|-------------|------|----------|-----------------|----------|
| | | | | |
| JP 02227079 | A2 | 19900910 | JP 1989-217540 | 19890825 |

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PRIORITY APPLN. INFO.:

JP 1988-250926 19881006

Human serum albumin fragments contg. the multiple sites for binding pharmaceuticals but not the C-terminal cysteine residues and the N-terminal long chain fatty acids binding sites are manufd. with recombinant Escherichia coli as carrier. Alternatively, the serum albumin fragments are manufd. as fusion products, esp. with the signal peptide of alk. phosphatase (phoA). Plasmid encoding a fusion protein of human serum albumin fragment (Met123-Pro303) and phoA signal peptide was constructed and transformed into E. coli. The E. coli transformants produced the fusion protein having a mol. wt. of 21000 (by SDSPAGE).
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             53 SEA FILE=REGISTRY SERUM ALBUMIN?/CN
L1
            330 SEA FILE=REGISTRY CHIMERIC
L3
             63 SEA FILE=REGISTRY ANGIOSTA?/CN
L4
             42 SEA FILE=REGISTRY ENDOSTATIN?
L5
            342 SEA FILE=REGISTRY CYSTINE?/CN
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             29 SEA FILE=REGISTRY TYROSINE KINASE?/CN AND RECEPTOR?
1.8
            209 SEA FILE=REGISTRY CYTOKINE?/CN AND RECEPTOR?
L9
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L11
            241 SEA FILE=REGISTRY ORPHAN?/CN
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          12190 SEA FILE=HCAPLUS CHIMERIC (5A) (PROTEIN? OR ?PEPTIDE?) OR L3
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            590 SEA FILE=HCAPLUS L4 OR ANGIOSTA?
L15
            346 SEA FILE=HCAPLUS L5 OR ENDOSTAT?
L16
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L18
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           1198 SEA FILE=HCAPLUS L13 AND (L15 OR L16 OR L17 OR L19)
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              9 SEA FILE=HCAPLUS L20 AND L14
L21
              9 SEA FILE=HCAPLUS L18 AND L20
L22
              6 SEA FILE=HCAPLUS L22 NOT L21
L23
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L23 ANSWER 1 OF 6 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2001:676999 HCAPLUS
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DOCUMENT NUMBER:

135:252790

TITLE:

Single nucleotide polymorphisms in human genes

INVENTOR(S):

Cargill, Michele; Ireland, James S.; Lander, Eric S.

PATENT ASSIGNEE(S):

Whitehead Institute for Biomedical Research, USA

SOURCE:

PCT Int. Appl., 145 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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APPLICATION NO. DATE
     PATENT NO.
                              DATE
                        KIND
                                                                  20010307
                                               WO 2001-US7268
                         A2
                               20010913
     WO 2001066800
          W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
              CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
          RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
              DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
              BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                            US 2000-187510
                                                             P 20000307
PRIORITY APPLN. INFO.:
                                                               P 20000522
                                            US 2000-206129
     The invention provides nucleic acid segments of the human genome,
AΒ
     particularly nucleic acid segments from genes including polymorphic sites.
     The polymorphisms were identified by resequencing of target sequences from
     individuals of diverse ethnic and geog. backgrounds by hybridization to
     probes immobilized to microfabricated arrays. Some of the single
     nucleotide polymorphisms (SNPs) specify a different amino acid sequence,
     some are silent or are in noncoding regions, and some specify a stop
     signal in protein translation. Allele-specific primers and probes
     hybridizing to regions flanking or contg. these sites are also provided.
     The nucleic acids, primers and probes are used in applications such as
     phenotype correlations, forensics, paternity testing, medicine and genetic
     147171-37-7, Adrenergic .beta. receptor kinase 2
ΙT
     149371-16-4, Adrenergic .beta. receptor kinase 1
     191941-10-3, prepronociceptin
     RL: BOC (Biological occurrence); BUU (Biological use, unclassified); PRP
      (Properties); BIOL (Biological study); OCCU (Occurrence); USES (Uses)
         (single nucleotide polymorphisms in human genes)
L23 ANSWER 2 OF 6 HCAPLUS COPYRIGHT 2002 ACS
                            2001:490351 HCAPLUS
ACCESSION NUMBER:
                            135:209360
DOCUMENT NUMBER:
                            Requirement for p38 and p44/p42 mitogen-activated
TITLE:
                            protein kinases in RAGE-mediated nuclear
                            factor-.kappa.B transcriptional activation and
                            cytokine secretion
                            Yeh, Chen-Hsiung; Sturgis, Lydia; Haidacher, Joe;
AUTHOR(S):
                            Zhang, Xue-Nong; Sherwood, Sidney J.; Bjercke, Robert
                            J.; Juhasz, Ondrej; Crow, Michael T.; Tilton, Ronald
                            G.; Denner, Larry
                            Department of Cell Biology and Apoptosis Program,
CORPORATE SOURCE:
                            Texas Biotechnology Corporation, Houston, TX, 77030,
                            Diabetes (2001), 50(6), 1495-1504
SOURCE:
                            CODEN: DIAEAZ; ISSN: 0012-1797
                            American Diabetes Association
PUBLISHER:
                            Journal
DOCUMENT TYPE:
LANGUAGE:
                            English
     Advanced glycation end product (AGE) activation of the signal-transducing
      receptor for AGE (RAGE) has been linked to a proinflammatory phenotypic
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change within cells. However, the precise intracellular signaling pathways involved have not been elucidated. We demonstrate here that human serum albumin modified with N.epsilon.-(carboxymethyl)lysine (CML), a major AGE adduct that progressively accumulates with aging, diabetes, and renal failure, induced nuclear factor (NF)-.kappa.B-driven reporter gene expression in human monocytic THP-1 cells. The NF-.kappa.B response was blocked with a synthetic peptide corresponding to the putative ligand-binding domain of RAGE, with anti-RAGE antiserum, and by coexpression of truncated receptors lacking the intracellular domain. Signal transduction from RAGE to NF-.kappa.B involved the generation of reactive oxygen species, since reporter gene expression was blocked with the antioxidant N-acetyl-L-cysteine. CML-modified albumin produced rapid transient activation of tyrosine phosphorylation, extracellular signal-regulated kinase 1 and 2, and p38 mitogen-activated protein kinase (MAPK), but not c-Jun NH2-terminal kinase. RAGE-mediated NF-.kappa.B activation was suppressed by the selective p38 MAPK inhibitor SB203580 and by coexpression of a kinase-dead p38 dominant-neg. mutant. Activation of NF-.kappa.B by CML-modified albumin increased secretion of proinflammatory cytokines (tumor necrosis factor-.alpha., interleukin-1.beta., and monocyte chemoattractant protein-1) severalfold, and inhibition of p38 MAPK blocked these increases. These results indicate that p38 MAPK activation mediates RAGE-induced NF-.kappa.B-dependent secretion of proinflammatory cytokines and suggest that accelerated inflammation may be a consequence of cellular activation induced by this receptor. THERE ARE 60 CITED REFERENCES AVAILABLE FOR THIS 60

L23 ANSWER 3 OF 6 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:136160 HCAPLUS

DOCUMENT NUMBER:

REFERENCE COUNT:

134:322241

TITLE:

Chemisorption of the Dipeptide Arg-Cys on a Gold Surface and the Selectivity of G-Protein Adsorption

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

Uvdal, K.; Vikinge, T. P.

AUTHOR(S): CORPORATE SOURCE:

Laboratory of Applied Physics Department of Physics

and Measurement Technology, Linkoeping University,

Linkoeping, S-581 83, Swed.

SOURCE:

Langmuir (2001), 17(6), 2008-2012 CODEN: LANGD5; ISSN: 0743-7463

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal English

LANGUAGE:

Arginine-L-cysteine dipeptide adsorbates are used in this study as a model system for G-protein-coupled receptors (GPCRs). An arginine-contg. model mol. is chosen

because the GPCR .alpha.2A has been shown to include an arginine-rich region in the G-protein-binding part of the third intracellular loop, and the role of arginines by means of recognition is believed to exceed their pos. charge. The dipeptide Arg-Cys is adsorbed to gold surfaces and the peptide monolayers are characterized. These peptide monolayers are then used for G-protein adsorption expts. to study the mol. interaction and binding. The mol. adsorption, orientation, and chem. binding of the peptide to the surface are studied by XPS and IR reflection-absorption spectroscopy. A chem. shift in the S(2p) core level spectrum of the

peptide adsorbate on gold shows that there is a strong mol. surface interaction consistent with a chem. binding of the peptide to the surface through the sulfur atom. With the cysteine part linked to the surface, the arginine part of the mol. is available for further adsorption processes. Monolayers of Arg-Cys, L-cysteine, and cysteamine are used for G-protein adsorption expts. Adsorption of human serum albumin and human Igs on the same monolayers are studied for comparison. The anal. tool is surface plasmon resonance. Two different buffers are used for the adsorption studies, and the influence of buffer compn. on protein adsorption is discussed. 32

REFERENCE COUNT:

THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L23 ANSWER 4 OF 6 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1999:795994 HCAPLUS

DOCUMENT NUMBER:

132:31744

TITLE:

Gene probes used for genetic profiling in healthcare

screening and planning

INVENTOR(S):

Roberts, Gareth Wyn

PATENT ASSIGNEE(S):

Genostic Pharma Ltd., UK PCT Int. Appl., 745 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PA' | PATENT NO. | | | | | DATE | | A | | CATIO | o | DATE | | | | | | |
|---------|-----------------------|-----|-----|-----|-----|------|-----|-----|------|-------|------|------|--------------|------|------|-----|-----|--|
| WO | O 9964627 A2 19991216 | | | | | | | | W | | | 0 | 19990604 | | | | | |
| | W: | AE, | AL, | AM, | ΑT, | AU, | ΑZ, | BA, | BB, | BG, | BR, | BY, | CA, | CH, | CN, | CU, | CZ, | |
| | | DE, | DK, | EE, | ES, | FI, | GB, | GD, | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, | |
| | | JP, | KE, | KG, | KP, | KR, | KZ, | LC, | LK, | LR, | LS, | LT, | LU, | LV, | MD, | MG, | MK, | |
| | | MN. | MW. | MX, | NO, | NZ, | PL, | PT, | RO, | RU, | SD, | SE, | SG, | SI, | SK, | SL, | ТJ, | |
| | | | | | | | | | | | | | | ΑZ, | | | | |
| | | | RU, | | | • | | | | | | | | | | | | |
| | RW: | | | | | MW, | SD, | SL, | SZ, | UG, | ZW, | ΑT, | ΒE, | CH, | CY, | DE, | DK, | |
| | | ES, | FI, | FR, | GB, | GR, | IE, | IT, | LU, | MC, | NL, | PT, | SE, | BF, | ВJ, | CF, | CG, | |
| | | CI, | CM, | GA, | GN, | GW, | ML, | MR, | ΝE, | SN, | TD, | TG | | | | | | |
| PRIORIT | Y APP | | | | | | | | | | 1209 | | Α | 1998 | 0606 | | | |
| | | | | | | | | | GB 1 | 998- | 1329 | 1 | Α | 1998 | 0620 | | | |
| | | | | | | | | | GB 1 | 998- | 1361 | 1 | Α | 1998 | 0624 | | | |
| | | | | | | | | | GB 1 | 998- | 1383 | 5 | Α | 1998 | 0627 | | | |
| | | | | | | | | | GB 1 | 998- | 1411 | 0 | Α | 1998 | 0701 | | | |
| | | | | | | | | | GB 1 | 998- | 1458 | 0 | Α | 1998 | 0707 | | | |
| | | | | | | | | | GB 1 | 998- | 1543 | 8 | Α | 1998 | 0716 | | | |
| | | | | | | | | | GB 1 | 998- | 1557 | 4 5 | A. | 1998 | 0718 | | | |
| | | | | | | | | | GB 1 | 998- | 1557 | 6 | Α | 1998 | 0718 | | | |
| | | | | | | | | | GB 1 | 998- | 1608 | 5 | Α | 1998 | 0724 | | | |
| | | | | | | | | | GB 1 | 998- | 1608 | 6 | \mathbf{A} | 1998 | 0724 | | | |
| | | | | | | | | | GB 1 | 998- | 1692 | 1 | Α | 1998 | 0805 | | | |
| | | | | | | | | | GB 1 | 998- | 1709 | 7 | Α | 1998 | 0807 | | | |
| | | | | | | | | | GB 1 | 998- | 1720 | 0 | Α | 1998 | 8080 | | | |
| | | | | | | | | | GB 1 | 998- | 1763 | 2 | Α | 1998 | 0814 | | | |

A 19980819 GB 1998-17943

There is considerable evidence that significant factor underlying the AB individual variability in response to disease, therapy and prognosis lies in a person's genetic make-up. There have been numerous examples relating that polymorphisms within a given gene can alter the functionality of the protein encoded by that gene thus leading to a variable physiol. response. In order to bring about the integration of genomics into medical practice and enable design and building of a technol. platform which will enable the everyday practice of mol. medicine a way must be invented for the DNA sequence data to be aligned with the identification of genes central to the induction, development, progression and outcome of disease or physiol. states of interest. According to the invention, the no. of genes and their configurations (mutations and polymorphisms) needed to be identified in order to provide crit. clin. information concerning individual prognosis is considerably less than the 100,000 thought to comprise the human genome. The identification of the identity of the core group of genes enables the invention of a design for genetic profiling technologies which comprises of the identification of the core group of genes and their sequence variants required to provide a broad base of clin. prognostic information - "genostics". The "Genostic.RTM." profiling of patients and persons will radically enhance the ability of clinicians, healthcare professionals and other parties to plan and manage healthcare provision and the targeting of appropriate healthcare resources to those deemed most in need. The use of this invention could also lead to a host of new applications for such profiling technologies, such as identification of persons with particular work or environment related risk, selection of applicants for employment, training or specific opportunities or for the enhancing of the planning and organization of health services, education services and social services.

9012-96-8, Cystathionase 54004-64-7, Rhodopsin kinase

127407-08-3, Receptor tyrosine kinase

138359-29-2, c-Kit protein tyrosine kinase

RL: ANT (Analyte); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(core group of disease-related genes; gene probes used for genetic profiling in healthcare screening and planning)

L23 ANSWER 5 OF 6 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1999:795993 HCAPLUS

DOCUMENT NUMBER:

132:31743

TITLE:

Gene probes used for genetic profiling in healthcare

screening and planning

INVENTOR(S):

Roberts, Gareth Wyn

PATENT ASSIGNEE(S): SOURCE:

Genostic Pharma Limited, UK PCT Int. Appl., 149 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

APPLICATION NO. DATE KIND DATE PATENT NO. 19990604 WO 1999-GB1779 WO 9964626 A2 19991216

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AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
              DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS,
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              MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ,
              TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ,
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                              19991230
                                               AU 1999-41586
                                                                  19990604
     AU 9941586
                         A1
                                               AU 1999-41587
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     GB 2339200
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         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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                                                              A 19980606
                                            GB 1998-12098
PRIORITY APPLN. INFO.:
                                            GB 1998-28289
                                                              A 19981223
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                                                              A 19980814
                                            GB 1998-17632
                                            GB 1998-17943
                                                              A 19980819
                                            WO 1999-GB1779
                                                               W 19990604
     There is considerable evidence that significant factor underlying the
AΒ
     individual variability in response to disease, therapy and prognosis lies
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in a person's genetic make-up. There have been numerous examples relating that polymorphisms within a given gene can alter the functionality of the protein encoded by that gene thus leading to a variable physiol. response. In order to bring about the integration of genomics into medical practice and enable design and building of a technol. platform which will enable the everyday practice of mol. medicine a way must be invented for the DNA sequence data to be aligned with the identification of genes central to the induction, development, progression and outcome of disease or physiol. states of interest. According to the invention, the no. of genes and their configurations (mutations and polymorphisms) needed to be identified in order to provide crit. clin. information concerning individual prognosis is considerably less than the 100,000 thought to comprise the human genome. The identification of the identity of the core group of genes enables the invention of a design for genetic profiling technologies.

9012-96-8, Cystathionase 54004-64-7, Rhodopsin kinase ΙT

127407-08-3, Receptor tyrosine kinase

138359-29-2, c-Kit protein tyrosine kinase

RL: ANT (Analyte); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(core group of disease-related genes; gene probes used for genetic profiling in healthcare screening and planning)

L23 ANSWER 6 OF 6 HCAPLUS COPYRIGHT 2002 ACS 1998:424365 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 129:91388

Recursive sequence recombination and screening as a TITLE:

tool for the in vitro evolution of gene products

INVENTOR(S): Patten, Phillip A.; Stemmer, Willem P. C.

PATENT ASSIGNEE(S): Maxygen, Inc., USA; Patten, Phillip A.; Stemmer,

Willem P. C.

SOURCE: PCT Int. Appl., 123 pp.

CODÉN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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KIND DATE
                                             APPLICATION NO.
    PATENT NO.
                             19980625
                                             WO 1997-US24239
                                                               19971217
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         W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
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             NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT,
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     EP 946755
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                                              JP 1998-528054
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     JP 2001506855
                                                                19990416
                             19990812
                                              AU 1999-23816
     AU 9923816
                        A1
                                          US 1996-769062
                                                           A1 19961218
PRIORITY APPLN. INFO .:
                                          AU 1995-29714
                                                             A3 19950217
                                          WO 1995-US2126
                                                            A2 19950217
                                          US 1995-564955
                                                            A2 19951130
                                          US 1996-537874
                                                            A2 19960304
                                                             A2 19960325
                                          US 1996-621859
                                                             A2 19960520
                                          US 1996-650400
                                          WO 1996-US19256 A2 19961202
                                          WO 1997-US24239 W, 19971217
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A method for development of proteins with new combinations of properties AB by recursive recombination of coding sequences of different origins and screening of gene products for desired properties is described. Recombination can be in vitro, or in vivo, e.g. using the cre/loxP system. Further variation can be introduced using mutagenesis-prone methods such as DNA repair. One method is denaturing and renaturing a population of fragments of 20-100 base pairs and selecting for those hybrids with base pair mismatches. These mismatched sequences are then ligated together to generate new sequences that will undergo DNA repair-mediated mutation. The method is flexible enough to allow coarse, or large scale, changes in sequences or it can be used at a very fine level: generating changes in a small subsequence. Many screening procedures may be used, but they must be carefully designed to detect changes of interest. Novel variants of calf intestinal alk. phosphatase with novel substrate specificity, human .alpha. interferon with higher specific activity, and luciferases with

increased stability are generated.

IT 86090-08-6, Angiostatin

RL: BSU (Biological study, unclassified); BIOL (Biological study) (development of novel variants of; recursive sequence recombination and screening as tool for in vitro evolution of gene products)

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=> d stat que
             53 SEA FILE=REGISTRY SERUM ALBUMIN?/CN
L1
L3
            330 SEA FILE=REGISTRY CHIMERIC
             63 SEA FILE=REGISTRY ANGIOSTA?/CN
L4
             42 SEA FILE=REGISTRY ENDOSTATIN?
L5
            342 SEA FILE=REGISTRY CYSTINE?/CN
L7
             29 SEA FILE=REGISTRY TYROSINE KINASE?/CN AND RECEPTOR?
L8
            209 SEA FILE=REGISTRY CYTOKINE?/CN AND RECEPTOR?
ь9
           2073 SEA FILE=REGISTRY G PROTEIN?/CN
L10
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L21
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L23
           1474 SEA FILE=HCAPLUS L17 AND LOOP?
L27
            107 SEA FILE=HCAPLUS L13 AND (L15 OR L16 OR L27 OR L19)
L28
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L29
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                ?CARCIN? OR ?NEOPLASM? OR ?SARCOM? OR ?LYMPHOM? OR ?MELANO? OR
                ?LEUKEM? OR ?METAST?)
             12 SEA FILE=HCAPLUS L29 NOT (L21 OR L23)
L30
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L30 ANSWER 1 OF 12 HCAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
                         2002:81559 HCAPLUS
TITLE:
                         Construction of eukaryotic expression vector of
                         endostatin gene and identification of its
                         activity
                         Fu, Luoan; Zhang, Xiang; Wu, Jingwen; Gao, Dakuan;
AUTHOR(S):
                         Yang, Lisun; Qu, Yan
                         Institute of Neurosurgery of Chinese PLA, Xijing
CORPORATE SOURCE:
                         Hospital, Fourth Military Medical University, Xi'an,
                         710033, Peop. Rep. China
                         Disi Junyi Daxue Xuebao (2001), 22(23), 2162-2165
SOURCE:
                         CODEN: DJDXEG; ISSN: 1000-2790
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Davis .09/768,183 Page 19

PUBLISHER:

Disi Junyi Daxue Xuebao Bianjibu '

DOCUMENT TYPE:

LANGUAGE:

Chinese

The expression vector of endostatin gene was constructed and its transfection into C6 cell and its antiangiogenesis activity were studied. The rat serum albumin secretive signal was linked to the endostatin gene C terminal by PCR, this fused gene was then inserted into polylinker sites of eukaryotic expression vector pcDNA3 to construct pcDNA-SE. The vector was transfected into C6 glioma cells by lipofectamine and the pos. clone was screened by G418. The activity of endostatin protein expressed by the C6 cells was examd. by immunohistochem. and the cell proliferation assay. The eukaryotic expression vector pcDNA-SE was successfully constructed and transfected into C6 cells. The cells expressed the endostatin protein which could inhibit the endotheliocyte proliferation. Endostatin was a potent angiogenesis inhibitor. The expt. led a foundation for following expts. on antiangiogenesis gene

L30 ANSWER 2 OF 12 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

therapy of tumors.

2002:74923 HCAPLUS

TITLE:

Improved survival in tumor-bearing SCID mice

treated with interferon-??-inducible protein 10

(IP-10/CXCL10)

AUTHOR(S):

Arenberg, Douglas A.; White, Eric S.; Burdick, Marie

D.; Strom, Scott R. B.; Strieter, Robert M.

CORPORATE SOURCE:

Department of Internal Medicine, Division of Pulmonary

and Critical Care Medicine, University of Michigan Medical School, Ann Arbor, MI, 48109-0642, USA Cancer Immunology Immunotherapy (2001), 50(10),

SOURCE: 533-538

CODEN: CIIMDN; ISSN: 0340-7004

PUBLISHER:

Springer-Verlag

DOCUMENT TYPE:

Journal

LANGUAGE:

AB

English

Tumor growth requires angiogenesis, which in turn requires an imbalance in the presence of angiogenic and angiostatic factors. We have shown that the CXC chemokine family, consisting of members that are either angiogenic or angiostatic, is a major determinant of tumor-derived angiogenesis in non-small-cell lung cancer (NSCLC). Intratumor injection of interferon-inducible protein 10 (IP-10, or CXCL10), an angiostatic CXC chemokine, led to reduced tumor growth in a SCID mouse model of NSCLC. In this study, we hypothesized that treatment with CXCL10 would, by restoring the angiostatic balance, improve long-term survival in NSCLC-bearing SCID mice. To test this hypothesis, A549 NSCLC cells were injected in the subcutis of the flank, followed by intratumor injections with CXCL10 continuously (group I), or for ten weeks (group II), or a control group (human serum albumin). Median survival was 169, 130, and 86 days resp. (P < 0.0001). We extended these studies to examine the mechanism of prolonged survival in CXCL10-treated mice. CXCL10 treatment inhibited lung metastases, but was dependent upon continued treatment, and was assocd. with an increased rate of apoptosis

in the primary tumor, with no direct effect on the proliferation of the NSCLC cells. Furthermore, the inhibition of lung metastases was due to the angiostatic effect of CXCL10 on the primary tumor, since the rate of apoptosis within lung metastases was unaffected. These data suggest that anti-angiogenic therapy of human lung cancer should be continued indefinitely to realize persistent benefit, and confirms the antimetastatic capacity of localized angiostatic therapy.

L30 ANSWER 3 OF 12 HCAPLUS COPYRIGHT 2002 ACS

2002:12702 HCAPLUS ACCESSION NUMBER:

Inhibition of spontaneous metastases TITLE:

formation by amifostine

Grdina, David J.; Kataoka, Yasushi; Murley, Jeffrey AUTHOR(S):

S.; Hunter, Nancy; Weichselbaum, Ralph R.; Milas, Luka

Department of Radiation and Cellular Oncology, CORPORATE SOURCE:

University of Chicago, Chicago, IL, 60637, USA

International Journal of Cancer (2002), 97(2), 135-141

CODEN: IJCNAW; ISSN: 0020-7136

Wiley-Liss, Inc. PUBLISHER:

Journal DOCUMENT TYPE: LANGUAGE:

SOURCE:

English Amifostine was investigated for its ability to inhibit spontaneous AΒ metastases formation using the well-characterized murine sarcoma, Sa-NH. Amifostine was administered i.p. at a dose of 50 mg/kg every other day for 6 days to C3Hf/Kam mice until tumors reached an av. size of 8-8.5 mm in diam. Amifostine was again administered immediately after surgical removal of the tumor -bearing limbs by amputation, and then once more 2 days later. Twenty-one days later, animals were evaluated for the presence of spontaneously developed pulmonary metastases. Nontumor-bearing control animals were sham treated using the same dosing and surgery schedules. Treatment with amifostine appeared to slightly delay tumor growth, i.e., 13 vs. 12 days for tumors to reach an av. diam. of 8 mm. Amifostine reduced both the incidence of pulmonary metastases formed in exptl. animals from 77% to 57% (p < 0.05), and their av. no. per animal from 12.8 .+-. 5.4 (SEM) to 2.9 .+-. 1.1 (SEM). The effect of amifostine exposure on serum levels of the angiogenesis inhibitor angiostatin was also detd. using Western blot anal. Consistent with the antimetastatic effect, exposure of animals to 50 mg/kg of amifostine resulted in a 4-fold enhanced serum level of angiostatin above control levels. phenomenon occurred in tumor-bearing and nontumor -bearing animals. The effects of amifostine on matrix metalloproteinase (MMP) enzymic activity was also detd. using gelatin zymog. Conditioned growth medium collected from sa-NH cells grown to confluency was exposed to various concns. of SH, i.e., 2-[(aminopropyl)amino]ethane-thiol (WR-1065), the active thiol form of amifostine, for either 30 min or 18 h. WR-1065, as a function of increasing dose and time, inhibited the enzymic activities of MMP-2 and MMP-9. At a concn. and time of exposure likely to be achieved in vivo, i.e., 40 .mu.M and 30 min, MMP-2 and MMP-9 activities were reduced to between 30% and 40% of control values. Consistent with these affects, WR-1065 was also found to be effective in inhibiting the ability of sa-NH cells to migrate through Matrigel membranes.

After an 18-h exposure under in vitro conditions, WR-1065 at concns. of 4, 40 and 400 .mu.M, and 4 mM, inhibited Sa-NH migration to 11%, 44%, 81% and 97% of control values, resp. The abilities of amifostine and its active thiol WR-1065 to stimulate angiostatin prodn. in mice, and to inhibit the MMP enzymic activities and invasion ability of Sa-NH cells under in vitro conditions, are consistent with the obsd. antimetastatic effects exhibited against Sa-NH tumors growing in vivo.

THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L30 ANSWER 4 OF 12 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:556856 HCAPLUS

DOCUMENT NUMBER:

135:286235

TITLE:

Genomic and proteomic analysis of the myeloid

differentiation program

AUTHOR(S):

Lian, Zheng; Wang, Le; Yamaga, Shigeru; Bonds, Wesley; Beazer-Barclay, Y.; Kluger, Yuval; Gerstein, Mark; Newburger, Peter E.; Berliner, Nancy; Weissman,

Sherman M.

CORPORATE SOURCE:

Department of Genetics, Boyer Center for Molecular Medicine, the Section of Hematology, Department of Internal Medicine, Yale University School of Medicine,

New Haven, CT, 06536-0812, USA Blood (2001), 98(3), 513-524 CODEN: BLOOAW; ISSN: 0006-4971 American Society of Hematology

PUBLISHER:

SOURCE:

Journal

DOCUMENT TYPE: English LANGUAGE:

Although the mature neutrophil is one of the better characterized AB mammalian cell types, the mechanisms of myeloid differentiation are incompletely understood at the mol. level. A mouse promyelocytic cell line (MPRO), derived from murine bone marrow cells and arrested developmentally by a dominant-neg. retinoic acid receptor, morphol. differentiates to mature neutrophils in the presence of 10.mu.M retinoic acid. An extensive catalog was prepd. of the gene expression changes that occur during morphol. maturation. To do this, 3'-end differential display, oligonucleotide chip array hybridization, and 2-dimensional protein electrophoresis were used. A large no. of genes whose mRNA levels are modulated during differentiation of MPRO cells were identified. results suggest the involvement of several transcription regulatory factors not previously implicated in this process, but they also emphasize the importance of events other than the prodn. of new transcription factors. Furthermore, gene expression patterns were compared at the level of mRNA and protein, and the correlation between 2 parameters was studied. 39 THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS

REFERENCE COUNT:

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L30 ANSWER 5 OF 12 HCAPLUS COPYRIGHT 2002 ACS 2001:473659 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

135:205729

TITLE:

Microarray analysis of the in vivo effects of hypophysectomy and growth hormone treatment on gene Davis .09/768,183 Page 22

Flores-Morales, Amilcar; Stahlberg, Nina; AUTHOR(S):

Tollet-Egnell, Petra; Lundeberg, Joakim; Malek, Renae

L.; Quackenbush, John; Lee, Norman H.; Norstedt,

Gunnar

Department of Molecular Medicine, Karolinska CORPORATE SOURCE:

Institute, Stockholm, 17176, Swed.

Endocrinology (2001), 142(7), 3163-3176 CODEN: ENDOAO; ISSN: 0013-7227 SOURCE:

Endocrine Society PUBLISHER:

DOCUMENT TYPE: Journal English LANGUAGE:

The authors used cDNA microarrays contg. 3000 different rat genes to study AB the consequences of severe hormonal deficiency (hypophysectomy) on the gene expression patterns in heart, liver, and kidney. Hybridization signals were seen from a majority of the arrayed cDNAs; nonetheless, tissue-specific expression patterns could be delineated. Hypophysectomy affected the expression of genes involved in a variety of cellular functions. Between 16-29% of the detected transcripts from each tissue changed expression level as a reaction to this condition. Chronic treatment of hypophysectomized animals with human GH also caused significant changes in gene expression patterns. The study confirms previous knowledge concerning certain gene expression changes in the above-mentioned situations and provides new information regarding hypophysectomy and chronic human GH effects in the rat. Furthermore, the authors have identified several new genes that respond to GH treatment. The results represent a first step toward a more global understanding of gene expression changes in states of hormonal deficiency.

92 THERE ARE 92 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT:

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L30 ANSWER 6 OF 12 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2001:315685 HCAPLUS

DOCUMENT NUMBER:

134:348950

TITLE:

Expression vector for human serum

albumin gene expression and its use in gene of

hypoalbuminaemia

INVENTOR(S):

Wood, Christopher Barry

PATENT ASSIGNEE(S):

UK

SOURCE:

Brit. UK Pat. Appl., 25 pp.

CODEN: BAXXDU .

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. KIND DATE _____ GB 1999-30891 19990805 GB 2350362 A1 20001129 GB 1998-17084 A 19980806 PRIORITY APPLN. INFO.:

This invention provides a vector construction comprising human

serum albumin gene which is used as gene therapy to treat the disorder of liver, like hypoalbuminemia. The plasmid vector pGT123 comprises myosin light chain 1/3 enhancer, CMV promoter and human serum albumin gene. The invention also provides the

pharmaceutical compn. of the gene therapy and examples of administration. Plasmid vector comprising human serum albumin gene was introduce into patient suffering the liver disorder which resulted in dramatic increase of the concn. of serum albumin in . blood.

ΙT 86090-08-6, Angiostatin 187888-07-9,

Endostatin

RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(gene for, in gene therapy of liver disease; vector construction for human serum albumin expression and use in gene therapy to treat hypoalbuminemia)

L30 ANSWER 7 OF 12 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2000:900985 HCAPLUS

DOCUMENT NUMBER:

134:126444

TITLE:

Expression of human angiostatin kringle(1-3)

in C6 cell suppresses endothelial cell

proliferation in vitro

AUTHOR(S):

Gao, Dakuan; Zhang, Xiang; Wu, Jingwen; Qu, Yan; Jing,

Junjie; Liang, Jingwen; Li, Shuhe

CORPORATE SOURCE:

Xijing Hospital, The Fourth Military Medical University, Xi'an, 710032, Peop. Rep. China Jiefangjun Yixue Zazhi (2000), 25(2), 83-86

SOURCE:

CODEN: CFCHBN; ISSN: 0577-7402

PUBLISHER:

Jenminjun Chubanshe

DOCUMENT TYPE:

Journal Chinese

LANGUAGE: To research anti-angiogenesis activity of, human AK(1-3)[

angiostatin kringle(1-3)], rat serum albumin

signal dipeptide was transfected into C6 cell utilizing lipofectamine. Electron microscope, flow cytometry, immunohistochem. and Western blot were used to detect the ultramicrostructure, cell cycle and expression of AK(1-3) of C6 cell transfected and untransfected. The antiangiogenesis activity of AK(1-3) expressed by C6 cell was identified with human umbilical vein endothelial cell proliferation assay. The results indicate that C6 cell successfully transfected with AK(1-3) gene can stably express active AK(1-3) which potently inhibits endothelial cell The study provides a promise for the antiproliferation. angiogenesis gene therapy in vivo.

L30 ANSWER 8 OF 12 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1999:325817 HCAPLUS

DOCUMENT NUMBER:

130:351218

TITLE:

Methods and compositions for enhancing immune response

and for the production of in vitro MABs Tamarkin, Lawrence; Paciotti, Giulio F.

INVENTOR(S):

Cytimmune Sciences, Inc., USA

PATENT ASSIGNEE(S):

PCT Int. Appl., 54 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

FAMILY ACC. NUM. COUNT:

English

PATENT INFORMATION:

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KIND DATE
                                                                    APPLICATION NO. DATE
       PATENT NO.
                                 ____
                                           _____
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                                            19990520
                                                                    WO 1998-US23957 19981110
                                    A2
       WO 9924066
       WO 9924066
                                    А3
                                            19991209
             W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
             W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                            19990531
                                                                 AU 1999-14548 · 19981110
       AU 9914548
                                  A1
                                                                    EP 1998-958518
                                                                                                19981110
                                    A2
                                            20001004
       EP 1039933
                  AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
                    IE, FI
                                                                     JP 2000-520153
       JP 2002503639
                                            20020205
                                                                                                19981110
                                    T2
                                                                US 1997-65155 P 19971110
PRIORITY APPLN. INFO.:
                                                                US 1998-75811
                                                                                         P 19980224
                                                                                         P 19981106
                                                                US 1998-107455
                                                                WO 1998-US23957 W 19981110
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The methods and compns. of the present invention are directed to enhancing AB an immune response and increasing vaccine efficacy through the simultaneous or sequential targeting of specific immune system components. More particularly, specific immune components, such as macrophages, dendritic cells, B cells and T cells, are individually activated by component-specific immunostimulating agents. One such component-specific immunostimulating agent is an antigen-specific, species-specific monoclonal antibody. The invention is also directed to a method for the in vitro prodn. of the antigen-specific, species-specific monoclonal antibodies which relies upon the in vitro conversion of blood-borne immune cells, such as macrophages and lymphocytes. Vaccine efficacy is enhanced by the administration of compns. contg. component-specific immunostimulating agents and other elements, such as antigens or carrier particles, such as colloidal methods, such as gold.

ΙT 86090-08-6, Angiostatin 187888-07-9, Endostatin

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (methods and compns. for enhancing immune response and for the prodn. of in vitro monoclonal antibodies)

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L30 ANSWER 9 OF 12 HCAPLUS COPYRIGHT 2002 ACS
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ACCESSION NUMBER:

1993:79450 HCAPLUS

DOCUMENT NUMBER:

118:79450

TITLE:

Kluyveromyces lactis as a host for the manufacture of

heterologous proteins

INVENTOR(S): PATENT ASSIGNEE(S): Fleer, Reinhard; Fournier, Alain; Yeh, Patrice

Rhone-Poulenc Rorer SA, Fr.

Eur. Pat. Appl., 16 pp. SOURCE:

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

French

Davis .09/768,183 Page 25

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

| | PAT | ENT I | .00 | | KIN | D | DATE | | | AI | PLIC | CATIO | ON NO | ٥. | DATE | |
|--------|--------------|--------------|-----|-------|----------|-----|--------------|------|-----|-------|-------|-------|-------|-----|------|------|
| | | 5217 | | | A1 | | 1993 | 0107 | | E | 2 199 | 92-40 | 01848 | в . | 1992 | 0630 |
| | FR | 2678 | 636 | | | | 1993 | | | FI | R 199 | 91-82 | 217 | | 1991 | 0702 |
| | | 2678 9301 | | | B1 A1 | | 1994 1993 | | | WC | 199 | 92-FI | R610 | | 1992 | 0630 |
| | | | | | | | JP, | | | | | | | | | ~= |
| | | RW: | AT, | BE, | CH, | | DK, | | | | | - | - | | NL, | |
| | CA | 2110 | | | AA | | | | | | | | | | 1992 | 0630 |
| | AU | 9222 | 783 | | A1 | | 1993 | 0211 | | Αl | J 199 | 92-22 | 2783 | | 1992 | 0630 |
| | ΕP | 5925 | 74 | | A1 | | 1994 | 0420 | | E | 199 | 92-93 | 15114 | 4 | 1992 | 0630 |
| | | R: | AT, | BE, | CH, | DE, | DK, | ES, | FR, | GB, | GR, | IT, | LI, | LU, | NL, | SE |
| | JΡ | 0650 | | | Т2 | | 1994 | 1020 | | JI | 199 | 92-50 | 02002 | 2 | 1992 | 0630 |
| | | 9304 | | | | | 1993 | | | | | | | | 1993 | |
| | US | 5633 | 146 | | Α | | 1997 | 0527 | | US | 199 | 95-45 | 54778 | 8 | 1995 | 0531 |
| PRIOF | | | | INFO. | : | | | | | FR 19 | 91-8 | 3217 | | | 1991 | 0702 |
| LICTOR | \ _ 1 | | | | • | | | | | US 19 | | | 17 | | 1992 | 0630 |
| | | | | | | | | | | WO 19 | | | | | 1992 | |
| | | | | | | | | | | | 1 | | - | | | |

An isolate of Kluyveromyces lactis CBS293.91, and mutants derived from it, are used to manuf. heterologous proteins. An expression vector for the manuf. of human serum albumin using the LAC4 promoter was constructed and introduced into a no. of isolates of K. lactis. K. lactis CBS293.91 showed lactose-inducible expression of the gene with yields of the protein considerably higher than when other strains of K. lactis were used. The generation of a mutant of K. lactis CBS293.91 analogous to the URA3 mutant of Saccharomyces cerevisiae is described. This mutant may be used for heterologous gene cloning/expression.

L30 ANSWER 10 OF 12 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1983:139125 HCAPLUS

DOCUMENT NUMBER:

98:139125

TITLE:

Biosynthesis of rabbit serum albumin

in a heterologous fractionated subcellular system Hradec, Jan; Stiborova, Marie; Dusek, Zdenek; Franek,

Frantisek

CORPORATE SOURCE:

Dep. Biochem., Oncol. Inst., Prague, CS-180 00/8,

Czech.

SOURCE:

Eur. J. Biochem. (1983), 131(2), 277-81

CODEN: EJBCAI; ISSN: 0014-2956

DOCUMENT TYPE:

Journal

LANGUAGE:

AUTHOR(S):

English

As demonstrated by a simple procedure based on indirect immunopptn., proteins retained on heparin-Sepharose 4B from postmitochondrial supernatants of rat liver and Zajdela hepatoma catalyze the translation of rabbit serum albumin mRNA in the presence of ribosomal subunits from rat liver, Zajdela hepatoma, or rabbit reticulocytes. The albumin synthesis shows an optimum at 1.5 mM MgCl2 and 25 mM KCl and requires ATP and GTP. It is significantly stimulated by tRNA and proceeds for >2 h, suggesting a high rate of reinitiation. At the optimum

ribosome:mRNA ratio of 13:1, the immunoprecipitable radioactivity was >15-20-fold the blank values. Fluorog. of polyacrylamide slabs after electrophoresis of immunoppts. revealed the presence of only complete full-size serum albumin without any smaller peptides resulting from premature terminations of polypeptide chains, demonstrating faithful translation. In stained gels only, both heavy and light chains of IgG were found, indicating that the assay procedure is highly specific and reliable. The fractionated heterologous protein -synthesizing system described in this paper may be generally useful for studies of the synthesis of specific proteins and factors affecting their rates since, unlike comparable translation assays, a precise calcn. of the balance of newly synthesized proteins is possible.

L30 ANSWER 11 OF 12 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1981:618698 HCAPLUS

DOCUMENT NUMBER: 95:218698

TITLE: Immunochemistry of conjugates prepared from serum

albumins and acridine nitrogen mustards (ICR mutagens)

AUTHOR(S): Creech, Hugh J.; O'Connell, Anna P.

CORPORATE SOURCE: Inst. Cancer Res., Fox Chase Cancer Cent.,

Philadelphia, PA, 19111, USA

SOURCE: Cancer Res. (1981), 41(10), 3844-51

CODEN: CNREA8; ISSN: 0008-5472

DOCUMENT TYPE: Journal

LANGUAGE: English

AB Antibodies elicited in rabbits by immunization with conjugates prepd. from serum albumins and nitrogen mustard derivs. of quinacrine (atebrin) had strong binding sites complementary to the quinacrine hapten. The characteristic absorption spectrum of quinacrine made possible accurate detns. of the antigen-antibody compn. of the serol. ppts. Conclusive evidence that such antibodies, in addn. to reacting with the quinacrine component of heterologous protein test conjugates, bind quinacrine itself, as well as closely related acridine haptens, was provided by quant. inhibition studies. Atebrin and the hydroxy precursors of several heterocyclic nitrogen mustards caused more than a 50% inhibition of the antigen-antibody reactions. The antibodies elicited by

of several heterocyclic nitrogen mustards caused more than a 50% inhibition of the antigen-antibody reactions. The antibodies elicited by the quinacrine-protein conjugates in ascites tumor-bearing mice substantially neutralized the antitumor effectiveness of the low dosages (0.5-2.0 .mu.mol/kg) of the acridine nitrogen mustards that were required for a demonstration of chemotherapeutic activity. In contrast, nitrogen mustard, which has no quinacrine moiety, was not affected. Immunization with unaltered serum albumin had no

influence on the activity of the acridine nitrogen mustards. Quant. in vitro inhibition studies allowed satisfactory predictions of in vivo immunol. reactivity.

L30 ANSWER 12 OF 12 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1956:53455 HCAPLUS

DOCUMENT NUMBER: 50:53455

ORIGINAL REFERENCE NO.: 50:10246h-i,10247a-b

TITLE: Immunological properties of carcinogen

-protein conjugates containing polycyclic aromatic

hydrocarbons and substituted stilbenes

AUTHOR(S): Creech, Hugh J.; Havas, H. Francis; Andre, Janet

-09/768,183Page 27 Davis

CORPORATE SOURCE:

Lankenau Hosp., Philadelphia

SOURCE:

Cancer Research (1955), 15, 726-33

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Antibodies toward 9,10-dimethyl-1,2-benzanthryl-3-carbamido-horse serum albumin conjugate reacted well with heterologous proteins contg. haptens which were either similar in size and configuration (3,4-benzopyrene) or smaller (1,2-benzanthracene, 2-acetylaminofluorene), but poorly with those contg. a larger structure (1,2,5,6-dibenzanthracene). The structural alteration produced by photooxidation of 9,10-dimethyl-1,2-benzanthracene was reflected by a change in haptenic activity. Conjugates in which 2'-methyl-4-dimethylaminostilbene was joined to protein by an azo or carbamido linkage showed haptenic activity and cross reactivity, but the 2 linkages were not equiv. The terminal dimethylaminophenyl portion of the substituted stilbene mol. was of serologic importance. After exhaustive sepn. of the antihapten-protein serum by native protein and by hapten-

heterologous protein, there remained much antibody which

was specifically pptd. by the completely homologous test antigen.